# **Refine Search**

### Search Results -

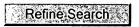
Terms	Documents
3\$1cyano\$12,4,5\$1trifluoro\$1benzoyl chloride	9

Database:

US Pre-Grant Publication Full-Text Database
US Patents Full-Text Database
US OCR Full-Text Database
EPO Abstracts Database
JPO Abstracts Database
Derwent World Patents Index
IBM Technical Disclosure Bulletins

Search:

5		 935
		(C)
		1









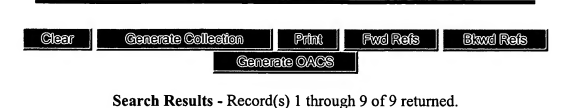
## Search History

DATE: Friday, September 02, 2005 Printable Copy Create Case

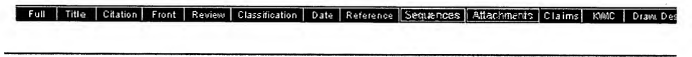
<u>Set Name</u> Query	•	Hit Count	Set Name
ide by side	•		result set
DB=USPT, USC	OC,EPAB,JPAB,DWPI,TDBD; PLUR=YES	S; OP=ADJ	
<u>L5</u> 3\$1cya	ano\$12,4,5\$1trifluoro\$1benzoyl chloride	9	<u>L5</u>
DB=USPT; PL	UR=YES; OP=ADJ		
L4 L3 sam	ne chlorinat\$7	13	<u>L4</u>
<u>L3</u> benzoy	ol chloride same benzoyl fluoride	71	<u>L3</u>
L2 L1 and	chlorinat\$7	44	<u>L2</u>
<u>L1</u> benzoy	ol chloride and benzoyl fluoride	107	<u>L1</u>

**END OF SEARCH HISTORY** 





- □ 1. Document ID: US 6706918 B2
- -: Invalid display element.



- ☐ 2. Document ID: US 6541675 B2
- -: Invalid display element.



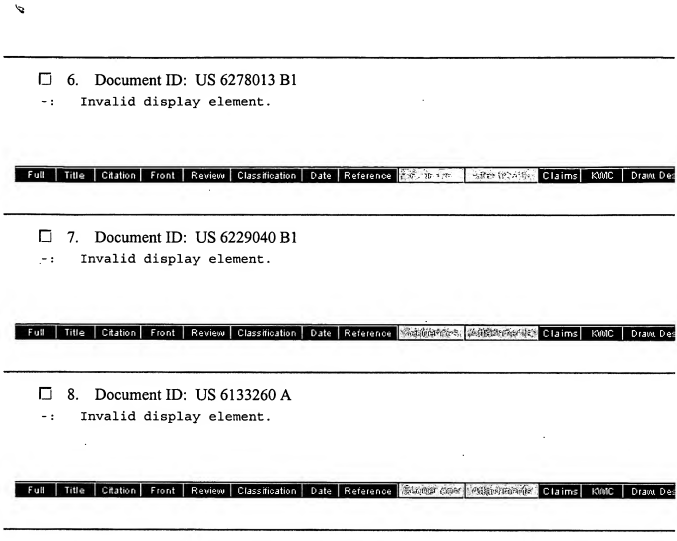
- ☐ 3. Document ID: US 6432948 B1
- -: Invalid display element.



- 4. Document ID: US 6323213 B1
- -: Invalid display element.

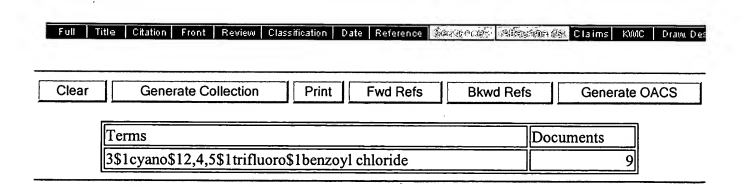


- □ 5. Document ID: US 6288081 B1
- -: Invalid display element.

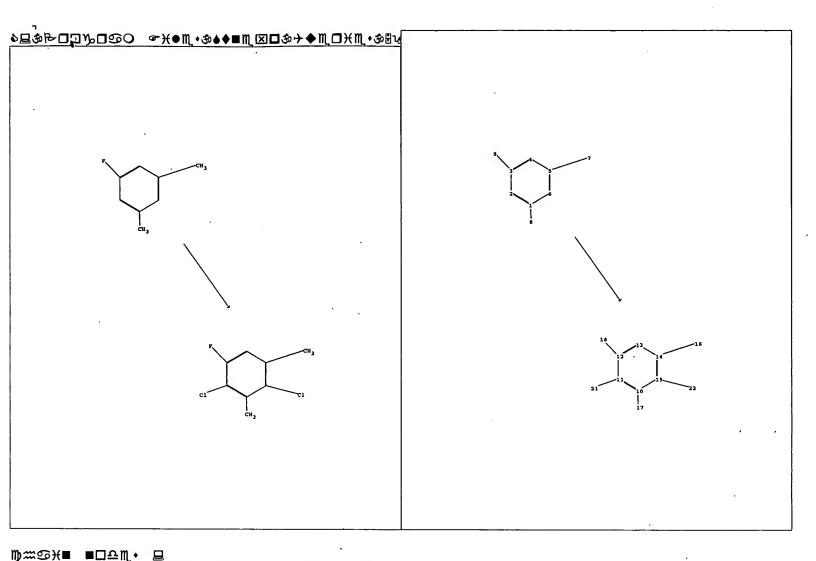


☐ 9. Document ID: CN 1119324 C, DE 19717231 A1, WO 9847862 A1, AU 9872163 A, EP 977729 A1, CN 1252789 A, HU 200003076 A2, US 6229040 B1, KR 2001005868 A, US 20010023300 A1, JP 2001521534 W, EP 977729 B1, DE 59803342 G, ES 2174431 T3, US 6541675 B2, US 20030092929 A1, HU 222056 B1, CN 1436771 A, IL 131974 A, US 6706918 B2, US 20040167350 A1

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Display Format: TI Change Format



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\* \* \* \* RECONNECTED TO STN INTERNATIONAL \* \* \* \* \* \* \* SESSION RESUMED IN FILE 'CAPLUS' AT 15:11:21 ON 02 SEP 2005 FILE 'CAPLUS' ENTERED AT 15:11:21 ON 02 SEP 2005 COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)

COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	60.02	499.01
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-7.30	-23.36
=> file casreact		
COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	60.02	499.01
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-7.30	-23.36

FILE 'CASREACT' ENTERED AT 15:11:33 ON 02 SEP 2005 USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications.

FILE CONTENT:1840 - 28 Aug 2005 VOL 143 ISS 9

New CAS Information Use Policies, enter HELP USAGETERMS for details.

Some CASREACT records are derived from the ZIC/VINITI database (1974-1991) provided by InfoChem, INPI data prior to 1986, and Biotransformations database compiled under the direction of Professor Dr. Klaus Kieslich.

This file contains CAS Registry Numbers for easy and accurate substance identification.

=>
Uploading C:\Program Files\Stnexp\Queries\593c.str

L15 STRUCTURE UPLOADED

=> d L15 HAS NO ANSWERS L15 STR

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

Structure attributes must be viewed using STN Express query preparation.

=> s 115
SAMPLE SEARCH INITIATED 15:12:26 FILE 'CASREACT'
SCREENING COMPLETE - 137 REACTIONS TO VERIFY FROM

2 DOCUMENTS

100.0% DONE 137 VERIFIED 0 HIT RXNS 0 DOCS SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

O SEA SSS SAM L15 (

BATCH \*\*COMPLETE\*\*

PROJECTED VERIFICATIONS:

2038 TO

3442 0

0 REACTIONS)

PROJECTED ANSWERS:

0 TO

=> s l15 sss full

FULL SEARCH INITIATED 15:12:41 FILE 'CASREACT'

SCREENING COMPLETE - 212 REACTIONS TO VERIFY FROM

20 DOCUMENTS

100.0% DONE

212 VERIFIED

0 HIT RXNS

0 DOCS

SEARCH TIME: 00.00.01

O SEA SSS FUL L15 ( O REACTIONS)

L17

L16

=>
Uploading C:\Program Files\Stnexp\Queries\593e.str

L23 STRUCTURE UPLOADED

=> d L23 HAS NO ANSWERS L23 ST

Structure attributes must be viewed using STN Express query preparation.

=> s 123

SAMPLE SEARCH INITIATED 15:33:03 FILE 'CASREACT'

SCREENING COMPLETE - 0 REACTIONS TO VERIFY FROM 0 DOCUMENTS

100.0% DONE 0 VERIFIED 0 HIT RXNS 0 DOCS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

BÂTCH \*\*COMPLETE\*\*

PROJECTED VERIFICATIONS: 0 TO 0

PROJECTED ANSWERS: 0 TO

L24 0 SEA SSS SAM L23 ( 0 REACTIONS)

=> s 123 sss full

FULL SEARCH INITIATED 15:33:14 FILE 'CASREACT'

SCREENING COMPLETE - 0 REACTIONS TO VERIFY FROM 0 DOCUMENTS

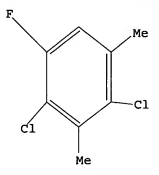
100.0% DONE 0 VERIFIED 0 HIT RXNS 0 DOCS

SEARCH TIME: 00.00.01

L25 0 SEA SSS FUL L23 ( 0 REACTIONS)

=>

=> d L26 HAS NO ANSWERS L26 STR



Structure attributes must be viewed using STN Express query preparation.

#### => s 126

#### REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

SAMPLE SEARCH INITIATED 15:37:47 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 38 TO ITERATE

100.0% PROCESSED 38 ITERATIONS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

BATCH \*\*COMPLETE\*\*

PROJECTED ITERATIONS: 391 TO 1129

PROJECTED ANSWERS: 0 TO 0

L27 0 SEA SSS SAM L26

L28 0 L27

=> s 126 full

#### REGISTRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

FULL SEARCH INITIATED 15:37:53 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 695 TO ITERATE

100.0% PROCESSED 695 ITERATIONS

1 ANSWERS

0 ANSWERS

SEARCH TIME: 00.00.01

1 SEA SSS FUL L26

L30 1 L29

L29

=> d ibib abs hitstr

L30 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1998:709043 CAPLUS

DOCUMENT NUMBER: 129:316044

TITLE: 3-Cyano-2,4,5-trifluorobenzoyl fluoride and

intermediates for its production INVENTOR(S): Marhold, Albrecht; Wolfrum, Peter PATENT ASSIGNEE(S): Bayer Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 30 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA'	TENT I	NO.	<b>.</b> - <b>.</b> -		KIN	<b>o</b> :	DATE			APF	LI	CAT	ION I	NO.			DATE	:	
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CA	2287	176			AA		1998	1029		CA	19	98-2	2287	176			1998	04	14
AU	9872	163			A1		1998	1113		ΑU	19	98-	7216	3			1998	04	14
EP	9777	29			A1		2000	0209		ΕP	19	98-	9192	66			1998	04	14
EP	9777	29			B1		2002	0313											
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I

GΙ

AΒ 3-Cyano-2,4,5-trifluorobenzoyl fluoride (I) is prepared starting from 5-fluoro-m-xylene and proceeding via 2,4-dichloro-5-fluoro-1,3dimethylbenzene, 2,4-dichloro-5-fluoro-3-(dichloromethyl)-1-(trichloromethyl) benzene, 2,4-dichloro-5-fluoro-3-(dichloromethyl) benzoic acid, 2,4-dichloro-5-fluoro-3-formylbenzoic acid (II), the oxime of II, and 2,4-dichloro-3-cyano-5-fluorobenzoyl chloride. 214774-61-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and chlorination of)

RN 214774-61-5 CAPLUS CN Benzene, 2,4-dichlor

Benzene, 2,4-dichloro-1-fluoro-3,5-dimethyl- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

### L18 🕆 STRUCTURE UPLOADED

=> d L18 HAS NO ANSWERS L18 STR

Structure attributes must be viewed using STN Express query preparation.

### => s 118

#### REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

SAMPLE SEARCH INITIATED 15:22:33 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 0 TO ITERAT

100.0% PROCESSED 0 ITERATIONS 0 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

BATCH \*\*COMPLETE\*\*

PROJECTED ITERATIONS: 0 TO 0 PROJECTED ANSWERS: 0 TO 0

L19 · 0 SEA SSS SAM L18

L20 0 L19

=> s 118 sss full

### REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

FULL SEARCH INITIATED 15:22:44 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 8 TO ITERATE

100.0% PROCESSED 8 ITERATIONS 1 ANSWERS

SEARCH TIME: 00.00.01

L21 1 SEA SSS FUL L18

### => d ibib abs hitstr

L22 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1998:709043 CAPLUS

DOCUMENT NUMBER: 129:316044

TITLE: 3-Cyano-2,4,5-trifluorobenzoyl fluoride and

> intermediates for its production Marhold, Albrecht; Wolfrum, Peter

INVENTOR(S): PATENT ASSIGNEE(S): Bayer Aktiengesellschaft, Germany SOURCE:

PCT Int. Appl., 30 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

P.	ATENT	NO.			KIN	)	DATE		I	APP:	LICAT	ION I	NO.		D	ATE	
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	6541				B2		2003										
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											1999-						
											2001-						
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AΒ 3-Cyano-2,4,5-trifluorobenzoyl fluoride (I) is prepared starting from 5-fluoro-m-xylene and proceeding via 2,4-dichloro-5-fluoro-1,3dimethylbenzene, 2,4-dichloro-5-fluoro-3-(dichloromethyl)-1(trichloromethyl)benzene, 2,4-dichloro-5-fluoro-3-(dichloromethyl)benzoic cid, 2,4-dichloro-5-fluoro-3-formylbenzoic acid (II), the oxime of II, and 2,4-dichloro-3-cyano-5-fluorobenzoyl chloride.

IT 214774-57-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and chlorination-dehydration of)

RN 214774-57-9 CAPLUS

CN Benzoic acid, 2,4-dichloro-5-fluoro-3-[(hydroxyimino)methyl]- (9CI) (CA INDEX NAME)

$$C1$$
 $CH \longrightarrow N-OH$ 
 $C1$ 

REFERENCE COUNT:

4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

Uploading C:\Program Files\Stnexp\Queries\593a.str

STRUCTURE UPLOADED

=> d L2 HAS NO ANSWERS

STR

Structure attributes must be viewed using STN Express query preparation.

=> s 12

#### REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

SAMPLE SEARCH INITIATED 14:52:34 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 99 TO ITERATE

100.0% PROCESSED 99 ITERATIONS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

BATCH \*\*COMPLETE\*\*

PROJECTED ITERATIONS: 1384 TO 2576

PROJECTED ANSWERS: 0 TO

L3 0 SEA SSS SAM L2

0 L3 L4

=> s 12 sss full

### REGISTRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

FULL SEARCH INITIATED 14:52:45 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 1932 TO ITERATE

100.0% PROCESSED 1932 ITERATIONS

0 ANSWERS

0 ANSWERS

SEARCH TIME: 00.00.01

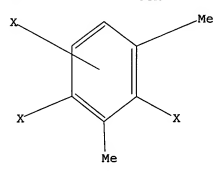
L6 0 L5

=>

#### STRUCTURE UPLOADED

=> d L7 HAS NO ANSWERS L7 STR

L7



Structure attributes must be viewed using STN Express query preparation.

### => s 17

### REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

SAMPLE SEARCH INITIATED 14:54:22 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 6255 TO ITERATE

32.0% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

BATCH \*\*COMPLETE\*\*

5 ANSWERS

PROJECTED ITERATIONS: 120359 TO 129841 PROJECTED ANSWERS: 75 TO 549

### => d 1-10 ibib abs hitstr

AUTHOR(S):

CORPORATE SOURCE:

L14 ANSWER 1 OF 10 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1991:428577 CAPLUS

DOCUMENT NUMBER: 115:28577

TITLE: Formation constants in carbon-hydrogen hydrogen

bonding. 2. Alkynes and polyhalobenzenes with

hexamethylphosphoric triamide in cyclohexane solution Lorand, John P.; Nelson, Jonathan P.; Gilman, R. Dean; Staley, Kathy L.; Chambers, Jeffery R.; Kirk, Harold

D.; Moeggenborg, Kevin J.; Farlow, David L.

Dep. Chem., Cent. Michigan Univ., Mt. Pleasant, MI,

48859, USA

SOURCE: Journal of Physical Organic Chemistry (1990

), 3(10), 659-69

CODEN: JPOCEE; ISSN: 0894-3230

DOCUMENT TYPE: Journal LANGUAGE: English

> Formation consts., K, for hydrogen bonding of weak C-H donors with hexamethylphosphoric triamide were measured in cyclohexane at 35° by PMR, data were analyzed by the Higuchi method. Donors include terminal alkynes and polyfluoro-, polychloro-, and polybromobenzenes. The K values for many of these are too small to measure in CCl4. For RC.tplbond.CH, K

decreases in the order R = BrCH2 > C6H5 > C2H5O > tert-Bu. For

polyhalobenzenes, K is generally larger for F than Cl or Br, and meta halogens facilitate hydrogen bonding most effectively.

Among tetrahalobenzenes, 1,2,3,4-isomers show the least evidence of 2:1 complex formation. Pentachlorobenzene has a smaller K than its

1,2,3,4-tetrachloro analog even after statistical correction; for C6HX5, K decreases in the order F > Cl > Br; the K values for

1,3,5-trihalobenzenes are closely similar; and 1,3,5-tribromo-2,4dimethylbenzene shows no evidence of hydrogen bonding by PMR. The last three observations suggest that buttressing is important in the C6HX5 structure, at least with C1 and Br. A value of K for benzene was measured, suggesting for the first time that the H atoms of benzene

engage in hydrogen bonding. The result compares well with values extrapolated from data for polyfluoro- but not polychlorobenzenes, again probably because of buttressing in the latter.

134636-45-6P, 1,3,5-Tribromo-2,4-dimethylbenzene

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and attempted hydrogen bonding of, to HMPT in cyclohexane)

RN 134636-45-6 CAPLUS

CN Benzene, 1,3,5-tribromo-2,4-dimethyl- (9CI) (CA INDEX NAME)

IT

L14 ANSWER 2 OF 10 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1988:473266 CAPLUS

DOCUMENT NUMBER: 109:73266

TITLE: New manganese tetrakis (halogenoaryl) porphyrins

featuring sterically hindering electronegative

substituents: synthesis of highly stable catalysts in

olefin epoxidation

AUTHOR (S): Banfi, Stefano; Montanari, Fernando; Quici, Silvio CORPORATE SOURCE:

Dip. Chim. Org. Ind., Univ. Milano, Milan, 20133,

Italy

SOURCE: Journal of Organic Chemistry (1988), 53(12),

2863-6

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal LANGUAGE: OTHER SOURCE(S): GI English CASREACT 109:73266

The complexes I (R = R2 = C1, Br, R1 = Me; R = R2 = H, R1 = C1) were prepared from halomesitylenes via condensation of halobenzaldehydes with pyrrole and complexation with Mn(OAc)2. I (R = C1, R1 = R2 = H; R = R2 = C1, Br, R1 = Me) are highly stable and efficient catalysts for the epoxidn. of cyclooctene and 1-dodecene with NaOCl at pH 9.5 and 0° under two-phase conditions, in the presence of N-hexylimidazole as axial ligand. The chemical stability dramatically decreases in I (R-R2 = H, F; R = R2 = Me, R1 = H; R = R2 = H, R1 = C1).

IT 608-72-0, 2,4,6-Tribromomesitylene

Ι

RL: RCT (Reactant); RACT (Reactant or reagent) (conversion to dimethyltribromobenzyl alc.)

RN 608-72-0 CAPLUS

CN Benzene, 1,3,5-tribromo-2,4,6-trimethyl- (9CI) (CA INDEX NAME)

CN Benzene, 1,3,5-trichloro-2,4,6-trimethyl- (9CI) (CA INDEX NAME)

IT 114634-36-5P 114634-37-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and complexation of, with manganese)

RN 114634-36-5 CAPLUS

CN 21H,23H-Porphine, 5,10,15,20-tetrakis(2,4,6-trichloro-3,5-dimethylphenyl)-(9CI) (CA INDEX NAME)

RN 114634-37-6 CAPLUS

CN 21H,23H-Porphine, 5,10,15,20-tetrakis(2,4,6-tribromo-3,5-dimethylphenyl)-(9CI) (CA INDEX NAME)

114634-34-3P 114634-35-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and coupling of, with pyrrole)

RN 114634-34-3 CAPLUS

CN Benzaldehyde, 2,4,6-tribromo-3,5-dimethyl- (9CI) (CA INDEX NAME)

IT

RN 114634-35-4 CAPLUS

CN Benzaldehyde, 2,4,6-trichloro-3,5-dimethyl- (9CI) (CA INDEX NAME)

IT

RN

114634-32-1P 114634-33-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and oxidation of)

RN 114634-32-1 CAPLUS

CN Benzenemethanol, 2,4,6-tribromo-3,5-dimethyl- (9CI) (CA INDEX NAME)

114634-33-2 CAPLUS

CN Benzenemethanol, 2,4,6-trichloro-3,5-dimethyl- (9CI) (CA INDEX NAME)

IT 114651-81-9P 114651-82-0P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of, as epoxidn. catalyst, stability of)

RN 114651-81-9 CAPLUS

CN Manganese, (acetato-0)[5,10,15,20-tetrakis(2,4,6-trichloro-3,5-

dimethylphenyl) -21H,23H-porphinato(2-)-N21,N22,N23,N24]-, (SP-5-12)- (9CI)
 (CA INDEX NAME)

RN 114651-82-0 CAPLUS

CN Manganese, (acetato-0) [5,10,15,20-tetrakis(2,4,6-tribromo-3,5-dimethylphenyl)-21H,23H-porphinato(2-)-N21,N22,N23,N24]-, (SP-5-12)- (9CI) (CA INDEX NAME)

L14 ANSWER 3 OF 10 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1977:422689 CAPLUS

DOCUMENT NUMBER:

87:22689

TITLE:

Permercurated arenes. Part II. Synthesis of perbromobenzoic acids and perbromobenzenes from aromatic carboxylic acids by permercuration and

bromodemercuration

AUTHOR (S):

Deacon, Glen B.; Farquharson, Graeme J.

CORPORATE SOURCE: Chem. Dep., Monash Univ., Clayton, Australia SOURCE:

Australian Journal of Chemistry (1977),

30(2), 293-303

CODEN: AJCHAS; ISSN: 0004-9425

DOCUMENT TYPE: Journal LANGUAGE: English

AB Mercuration of RC6H4CO2H (R = 2-Me, -Cl, -F, and -Br,3-Me, -F, -C1, -Br, -CF3, -NO2, and -OMe, and 4-Me, -F, -C1, -Br, -CF3, and -NO2) and 2,6-R2C6H3CO2H (R = Me,

C1, Br) with (CF3CO2)2Hg and subsequent bromodemercuration gave

the resp. RC6Br4CO2H and 2,6-R2C6Br3CO2H, which were accompanied by C6Br5R

and 1,3-R2C6Br4. 39568-70-2P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 39568-70-2 CAPLUS

CN Benzene, 1,2,3,5-tetrabromo-4,6-dimethyl- (9CI) (CA INDEX NAME)

IT

L14 ANSWER 4 OF 10 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1976:179885 CAPLUS

DOCUMENT NUMBER: 84:179885

TITLE: Quality of phthalic acids improved by haloacetic acid

INVENTOR(S): Kuhlmann, George E. PATENT ASSIGNEE(S): Standard Oil Co., USA

SOURCE: U.S., 6 pp.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 3947494	Α	19760330	US 1972-316857	19721220 <
PRIORITY APPLN. INFO.:			US 1972-316857	A 19721220
AB Phthalic acids of	improved	quality we	re obtained by direct	t oxidation of
			d p-Me2C6Br4, p-Me2C	
an oxidation zone	with sma	ll amts. of	haloacetic acid, e.g	g., XCH2CO2H (X =
<b>Cl</b> , Br, <b>F</b> ) or F3C	CO2H, pre	sent in HOA	c solution of one or	
more heavy metal	oxidation	catalysts	and Br-releasing Br-	containing compound
BrCH2CO2H was the				3 1
TT 20560 70 0				

IT

RL: RCT (Reactant); RACT (Reactant or reagent) (oxidation of, effect of haloacetic acids on quality of phthalic acids

RN 39568-70-2 CAPLUS

CN Benzene, 1,2,3,5-tetrabromo-4,6-dimethyl- (9CI) (CA INDEX NAME)

L14 ANSWER 5 OF 10 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1972:447779 CAPLUS

DOCUMENT NUMBER: 77:47779

TITLE: Stable carbocations. CXXXIX. Nitro- and

chlorohexamethylbenzeneium ions and 1-nitro- and

1-chloro-2,4,6-trifluoromesitylenium ions Olah, George A.; Lin, Henry C.; Mo, Y. K.

CORPORATE SOURCE: Dep. Chem., Case West. Reserve Univ., Cleveland, OH,

USA

SOURCE: Journal of the American Chemical Society (1972

), 94(10), 3667-9

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Journal LANGUAGE: English

GI For diagram(s), see printed CA Issue.

Hexamethylbenzeniums (I, X = NO2, Cl) and 2,4,6-trifluoro-mesityleniums (II, X = NO2, Cl) were obtained by treating Me6C6 or 1,3,5,2,4,6-F3Me3C6 with NO2+BF4-FSO3H-SO2 and SbF5-FSO3H-Cl2-SO2ClF solution at low temperature. The structure of the ions were confirmed by PMR and F NMR. The PMR of I (R = NO2) is temperature dependent indicating the degenerate migration or exchange process of the nitro group (NO2+). The energy of activation of the process was 16.8 ± 1.5 kcal/mole.

IT 363-64-4

AUTHOR (S):

AB

RN

RL: PRP (Properties)

(benzenium ion formation from)

363-64-4 CAPLUS

CN Benzene, 1,3,5-trifluoro-2,4,6-trimethyl- (9CI) (CA INDEX NAME)

L14 ANSWER 6 OF 10 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1963:441438 CAPLUS

DOCUMENT NUMBER: 59:41438

ORIGINAL REFERENCE NO.: 59:7428d-h,7429a-c

TITLE: Perchloryl aromatic compounds

INVENTOR(S): Inman, Charles E.; Oesterling, Robert E.; Tyezkowski,

Edward A.

PATENT ASSIGNEE(S): Pennsalt Chemicals Corp.

SOURCE: 12 pp.
DOCUMENT TYPE: Patent

LANGUAGE: Unavailable

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

US 3076853 19630205 US 19620621 <-The title compds. ArClO3 were prepared by treating the appropriate aromatic

The title compds. ArClO3 were prepared by treating the appropriate aromatic compound with ClO3F in the presence of a Freidel-Crafts catalyst. AlCl3 was the preferred catalyst. An excess of the aromatic compound or an inert solvent may be used. Thus, AlCl3 133 was suspended with agitation in benzene 2600 parts in a cooling bath. ClO3F gas 100 patts was slowly passed into the mixture and the temperature maintained at 40°. Gaseous HCl was evolved. After HCl evolution had ceased, the reaction mass was added to twice its volume of H2O, and steam-distilled The benzene layer was evaporated and distilled in vacuo to give perchlorylbenzene 62 parts, b. 232°, f.p. -3°, n2DO 1.5236, d3O 1.185. Similarly prepared were the following substituted perchlorylbenzenes [substituent(s) given]:

2,4-dimethyl; 2,5-dimethyl (b2 78°, m. 27-8°); 4-fluoro

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(b0,25 53°,n2D0 1.5051); 2,5dibromo; 2,5-dimethyl; 2,5-difluoro;
3,4-dimethyl; 4-hydroxy; 3,4-dibromo; 2,4-dimethyl; 2,5-dimethyl-4-chloro;
3,4,5-trichloro; 2,5-dichloro-3-fluoro; 2,4,6-trimethyl;
2,3,5,6-tetramethyl; 2-chloro-5-fluoro. A solution of perchlorylbenzene 20
in concentrated H2SO480 was treated with a mixture of concentrated H2SO425 and concentrated
HNO3 14 parts at 20-300 for 1 hr. The mixture was poured over ice and a
yellow solid 22 parts was filtered from the mixture Recrystn. from
benzene-petr. ether gare yellow 3-nitroperchlorylbenzene, m.
49-50°, reduced with SnCl2 in alc. HCl to 3-aminoperchlorylbenzene,
m. 32°. The following substituted perchlorylbenzenes were prepared
by known methods [substituent(s) given]: B-NH2.HCl; 3-acetamido (m. 136
70); 4-C1; 4-Me; 3,4-dichloro; 3-amino-4-methyl; 3- amino- 4-
methyl- 6- nitro; 2,3,5-trichloro-4-fluoro; 2,3,5,6-tetrachloro-4-hydroxy-;
 3-HO3S; 2-MeO; 2-Cl; 2-chloro-5-nitro; 4-MeS;
3-(4-nitrophenyl); 4nitro-3-(4-nitrophenyl); 2-chloro-5-amino;
2-chloro-3-nitro-5amino; 2-Me; 3,5,6-trichloro-2-methyl;
3,4,5-trichloro-2-methyl; 2,3,4,5,6-pentachloro; 2,5-dimethyl-3-cyano;
3,4-dichloro-2,5difluoro; 3,6-dichloro-2,5-difluoro; 3,4-dichloro-5-
methoxy; 3,4dichloro-5-phenyl; 3,4-dibromo-2,5-dichloro;
2,4-bis(chloromethyl); 2,4,5-trimethyl-3-nitro; 3,4,5-trichloro-2-iodo;
2,4,5,6-tetrachloro-3-fluoro: 3-iodo-2,4,6-trimethyl; 4-amino-
2,3,5,6tetramethyl; 3-acetamido-2-nitro; 2,5-diamino-3-chloro;
3(3-perchlorylphenyl); 4-chloromethyl. Also prepared were
azobis(3-perchlorylbenzene), azoxybis(3-perchlorylbenzene), and
hydrazobis(3-perchlorylbenzene). 3-Aminoperchlorylbenzene-HCl 1 part was
dissolved in 20% HCl and diazotized with NaNO2 solution at 0-5° to
form a solution of the diazonium salt. Yellow crystals of
3,3'-diperchloryldiazoaminobenzene were removed by filtration. To the
filtrate was added \beta-naphthol 1 part dissolved in dilute NaOH to give
dark orange 3-(\beta-hydroxynaphthylazo)perchlorylbenzene, m.
200-1°. A mixture of 0.0233 mole 4-fluoroperchlorylbenzene and 0.221
mole NaOMe in 200 ml. MeOH was refluxed for 90 min., the cooled mixture
diluted with H2O, and extracted with Et25 to give 4-methoxyperchlorylbenzene,
n2D0 1.5307. 4-Phenylthioperchlorylbenzene, n2D6 1.5778, was similarly
prepared from 4-finoroperchlorylbenzene and thiophenol in the presence of
Na-OMe. 4-Chloroperchlorylbenzene was also prepared by chlorination of
perchtorylbenzene at 10-16° in the presence of 1% anhydrous FeCl3.
Other nuclear chlorinated compds. were similarly prepared by continuing
chtorination below 280° to substitution tip to 5 Cl
       Bromo- and iodoperchlorylbenzene compds. were simi- larly prepared
using the appropriate halide catalyst (FeBr3, iodine, or Fe).
Alkylhaloperchloryl aryl compds. were prepared by halogenating the
alkylperchloryl aryl compound Thus, 2,4-dimethyl-perchlorylbenzene was
progressively chlorinated, first at 5-10° and then by gradually
raising the temperature to 60°, with gaseous Cl in the presence
of FeCl3 and in a suitable solvent (ClCH2CCL3, CCL14, PhNO2, Et20, EtOH,
CHCl3, or glacial HO3c) to give 3-chloro-, 3,5-dichloro-, and
3,5,6-trichloro-2,4-dimethylperchloryl-benzene. Chlorination of
2,4-dimethylperchlorylbenzene was carried out in a glass tower packed with
glass rings and illuminated with Hg lamps spaced 4 ft. apart.
2,4-dimethyl-perchlorylbenzene was heated to 65-75° and red into
the top of the tower at a uniform rate. Dry Cl gas was passed
up the tower. The temperature of the tower was kept just below the reflux point
and 2,4-chloromethylperchlorylbenzene was recovered as product. Highly
alkylated and hydroxy-containing derivs. of perchloryl-benzene were
chloromethylated by treating the perchloryl aryl compound with HCHO and HCl
below 280° in the presence of H2SO4, ZnCl2, Al, or Sn as catalysts.
Perchlorytbenzene and its homologs were sulfonated by heating a mixture of
the perchtoryl aryl compound and concentrated H2SO4 below 280°. The title
compds. were useful as explosives. 3-(\beta-Hydroxynaphthylazo)perchloryl-
benzene was useful as a dye for wool.
90048-78-5, m-Xylene, 2,4,5-trichloro-6-perchloryl-
90151-28-3, m-Xylene, 2,4-dichloro-6-perchloryl-
   (preparation of)
90048-78-5 CAPLUS
m-Xylene, 2,4,5-trichloro-6-perchloryl- (7CI) (CA INDEX NAME)
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IT

RN

CN

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RN90151-28-3 CAPLUS

m-Xylene, 2,4-dichloro-6-perchloryl- (7CI) (CA INDEX NAME)

L14 ANSWER 7 OF 10 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1963:81229 CAPLUS

DOCUMENT NUMBER: 58:81229 ORIGINAL REFERENCE NO.: 58:13841b-h

TITLE: Perchloryl aromatic compounds

INVENTOR (S): Inman, Charles E.; Oesterling, Robert E.; Tyezkowski,

Edward A.

PATENT ASSIGNEE(S): Pennsalt Chemicals Corp.

SOURCE: 12 pp. DOCUMENT TYPE: Patent LANGUAGE: Unavailable

PATENT INFORMATION:

AB

PATENT NO. KIND DATE APPLICATION NO. DATE ---------US 3070609 19621225 US 19620621 <--

The F atom of FClO3 (I) (perchloryl fluoride, b. -47.5°), was replaced with a substituted or unsubstituted aryl radical, by using a new organic unit process, perchlorylation (similarly to nitration, sulfonation, etc.), to form perchloryl aromatic compds. These can be further treated (sulfonated, halogenated, etc.). The new compds. possess explosive properties; they are useful as explosive charges for blasting and for the manufacture of explosive devices, also as high energy fuels, as intermediates in the preparation of compds. useful in pharmaceutical and dye application, as additives for fuels in internal combustion engines, particularly as cetane improvers in diesel fuels. To prepare these compds., I in gaseous form and the aromatic compound (preferably C6H6, or substituted C6H6 as PhMe, PhEt, xylenes, mono-, di-, and trihalobenzenes; and chlorotoluenes) are brought together in stoichiometric proportions in the presence of Friedel-Crafts catalysts, as AtCl3, about mole per mole, in a nonalk., nonaq. system at 0-80° (preferably 0-15° for most of the reactions). Preferably, the catalyst is added to the aromatic compound, with the latter being used alone in excess or dissolved in a nonaq., inert solvent or diluent, and I is then passed into the mixture with agitation. The perchlorylated aromatic compds. can be treated under mildly alkaline, neutral, or acid conditions, below 280% around which temperature the ClO3 group is unstable and the compds. decompose explosively. The ClO3 group is replaced by HO group upon treatment with inorg. or organic base under strongly alkaline conditions. Thus, 133 parts AlCl3 was suspended with agitation in 2600 parts C6H6 in a vessel in a cooling bath. I gas was passed slowly into the C6H6-AlCl3 mixture which was maintained at 40°. HCl gas was evolved from the reaction mass. Addition of I was

stopped when 100 parts had been added and HCl evolution ceased. reaction mass was added to twice its volume of water. The mass was then steam-distilled The C6H6 layer thus recovered was evaporated, and the PhClO3 contained therein, 70 parts, was recovered as a pale yellow oil. Distillation in vacuo gave 62 parts PhClO3, b. 282° f.p. -3° n20D 1.5236, d30 1.186. Similarly, from m-C6H4(CH3)2 (with AlCl3 and I), 2,4-Me2C6H3ClO3 was obtained, b. 78°. From PhF (with AlCl2 and I), 4-FC6H4 ClO2was obtained, b0.25 53°, n20D 1.5051. From 3-02NC6H4ClO3 (with EtOH, concentrated HCl, SnCl2), 3-H2NC6H4ClO3 was obtained, m.32°. From PhClO3 (with concentrated H2SO4, concentrated HNO3), 3-O2-NC6H4ClO3 was obtained, m. 49-50°. O2NC6H4Ph (with I and AlCl3) gave perehlorylnitrobiphenyl. From PhOH (with I and AlCl3), 4-HOC6H4ClO3 was obtained. From 3-H2NC6H4ClO3 (with anhydrous ether, anhydrous HCl), the HCl salt was obtained, decomposed From 3-H2NC4H4ClO3 (with Ac20-AcOH), 3-AcNH-C6H4ClO3 was obtained, m. 136-7° (EtOH). PhClO3 at atmospheric pressure and 285° detonated vigorously; in liquid and in solid form it exploded when it was subjected to impact on a detonating block. m-O2-NC6H4ClO3, subjected to impact on a detonating block, exploded, p-02NC6H4Cl03 was exploded upon detonation of blasting caps by means of an elec. detonator. 3-H2NC6H4ClO3.HCl (with 20% HCl, diazotized with NaNO2 solution) gave 3-O3ClC6H4N: -N NHC6H4ClO3-3, which with  $\beta$ -naphthol in dilute NaOH gave a dye, 3-( $\beta$ -HOC10H6N:N)C6H4ClO3, dark orange crystals, m. 200-1°, λ 464 mμ. Wool cloth was dyed with 3-(β-HOC10H6N:N)C6H4-CIO3 to a deep orange color, which withstood prolonged exposure in the sunlight without appreciable loss of 4-FC6H4ClO3 with MeONa-MeOH gave 4-MeOC6H4ClO3, n28D 1.5307. 4-F-C6H4ClO3 with HSPh-MeONa gave 4-PhSC6H4ClO3, n26D 1.5778. From PhClO3 (with FeCl3 and gaseous C1), 4-CLC6H4ClO3, was obtained, all oily liquid. 90048-78-5, m-Xylene, 2,4,5-trichloro-6-perchloryl-

IT 90151-28-3, m-Xylene, 2,4-dichloro-6-perchloryl-(preparation of) RN

90048-78-5 CAPLUS

CN m-Xylene, 2,4,5-trichloro-6-perchloryl- (7CI) (CA INDEX NAME)

RN90151-28-3 CAPLUS CN m-Xylene, 2,4-dichloro-6-perchloryl- (7CI) (CA INDEX NAME)

L14 ANSWER 8 OF 10 CAPLUS COPYRIGHT 2005 ACS on STN

1955:45417 ACCESSION NUMBER: CAPLUS

DOCUMENT NUMBER: 49:45417 ORIGINAL REFERENCE NO.: 49:8700e-f

TITLE: Absorption spectra in relation to the chemical reactivity of some haloaromatic compounds

AUTHOR (S): Smith, F.; Turton, Lilian M.

CORPORATE SOURCE: Univ. Birmingham, UK

SOURCE: Journal of the Chemical Society, Abstracts (

1955) 1350-4

CODEN: JCSAAZ; ISSN: 0590-9791

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

AB cf. J. Appl. Chemical 4, 325(1954). The ultraviolet-light-absorption spectra

of a number of C1- and F-aromatic compds. are described,

and some correlation of the results with the degree of halogenation of the substances and their reactivity is attempted. The spectrophotometric measurements were carried out as previously described (cf. C.A. 45,

8352d).

IT

**5324-68-5**, Mesitylene, 2,4,6-trichloro-

(spectrum of)

RN5324-68-5 CAPLUS

CN Benzene, 1,3,5-trichloro-2,4,6-trimethyl- (9CI) (CA INDEX NAME)

L14 ANSWER 9 OF 10 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1953:11438 CAPLUS

DOCUMENT NUMBER: 47:11438

ORIGINAL REFERENCE NO.: 47:2036i,2037a-e

Electron distribution in molecules. I. F19 nuclear

magnetic shielding and substituent effects in some

benzene derivatives

AUTHOR (S): Gutowsky, H. S.; McCall, D. W.; McGarvey, B. R.;

Meyer, L. H.

CORPORATE SOURCE: Univ. of Illinois, Urbana

SOURCE: Journal of the American Chemical Society (1952

), 74, 4809-17

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

The effect of substituents on the electron distribution in benzene produces changes in the nuclear magnetic shielding of F atoms in the mol. A comparison of the F19 nuclear magnetic shielding in PhF with that in a substituted PhF was used in defining a  $\delta$ -parameter (C.A. 46, 5433b). A radiofrequency spectroscope with slow sweep modulation was used to measure mol. differences in nuclear magnetic shielding. Exptl. δ-values were determined for the following monosubstituted PhF derivs.: o-, m-, and p-NO2, -CN, -CO2H, -I, -Br, -C1, -F, -Me, -NHAc, -OH, -OEt, -NH2; o- and p-OMc; p-SO2Cl, -CCl3, -Ph, -OC6H4F, -NHPh, -NMe2; o-CH:CHCO2H; m-CF3, -CH(OH)Me, -OC6H4F. A linear correlation was

observed between Hammett's substituent constant  $\sigma$  and measured  $\delta$ 

values. A least-squares solution for meta derivs. gives σm =

1.698m and for para derivs.,  $\sigma p = 0.5608p + 0.271$ .

Systematic differences in the correlation were observed and attributed to the dependence of the  $\delta$ -values on the nature of the electronic

interactions of the substituent. Thereby, a detailed analysis of the

 $\delta\text{-}$  and  $\sigma\text{-}values$  for particular substituents permits evaluation of the nature of their electronic effects. Consideration of

o-substituents suggests that an interaction contributes to the

So-values in some cases in addition to the usual inductive and electromeric effects at the m- and p-positions. The potential uses and

limitations of δo-values in evaluating o-effects and entropies of

reaction are discussed. A determination of the extent to which substituent effects are additive was made by measuring  $\delta$ -values for the

```
following polysubstituted PhF compds. (observed F at the
1-position): 3-NO2, 4-NH2; 2-NO2, 4-NH2; 3, 5-, 2, 4-, 3, 4-, and 2,
5-di-fluoro; 3-F, 5-1; 2, 4-dinitro; 2-NH2, 4-F;
3-NH2, 4-F; 2-NH2, 5-F; 3-F, 4-NH2; 2-NO2,
4-F; 3-NO2, 4-F; 2-NO2, 5-F; 3-F,
4-NO2; 2-Cl, 4-F; 3-Cl, 4-F; 2-
Cl, 5-F; 3-F, 4-Cl; 2-Br, 4-
F; 3-Br, 4-F; 2-I, 4-F; 3-I, 4-F;
2-I, 5-F; 3-F, 4-I; 2-CF3, 4-F; 3-CF3, 4-
F; 2,4,6-trimethyl; 2,5-dichloro-4-F; 2, 5-dibromo-4-
F; 2,4,5-trifluoro; 2, Br-4,5-difluoro; 2,4-difluoro-5-Br;
2,5-difluoro-4-Br; 2,5-difluoro-3-CF3; 2,4-difluoro-6-CF3;
3,4-difluoro-5-CF3; 2,4,6-trimethyl-3-F; 2,4,6-trimethyl-3,5-
difluoro. The electronic effects of substituents are usually additive,
but there is significant nonadditivity between substituents in about 1/3
of the compds. studied, demonstrating interaction between substituents. A
\delta' parameter, analogous to \delta, is defined in terms of the
change produced by a substituent in the ring on the F19 nuclear magnetic
shielding in benzotrifluorides. The \delta'-values were determined for the
following substituted benzotrifluorides: o-, m-, and p-Cl, -CF3;
o- and m-NH2; m-F, -Br, -NO2. The \delta'-values are about
1/10 of the \delta-values for the same substituents; this indicates the
extent of attenuation of the substituent effects in the side chain.
363-64-4, Mesitylene, 2,4,6-trifluoro-
   (fluorine nuclear magnetic shielding in)
363-64-4 CAPLUS
Benzene, 1,3,5-trifluoro-2,4,6-trimethyl- (9CI)
                                                  (CA INDEX NAME)
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ACCESSION NUMBER:

CORPORATE SOURCE:

ORIGINAL REFERENCE NO.:

DOCUMENT NUMBER:

TITLE:

SOURCE:

AUTHOR (S):

IT

RN

CN

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(Washington, D. C.) (1947), 39, 64-8
                         CODEN: JIECAD; ISSN: 0095-9014
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         Unavailable
AB
     In a search for new types of compds. which vulcanize GR-S without S with
     formation of products with superior resistance to heat, a new class of
     vulcanizing agents was found comprising halogenated organic compds. which can
     in turn be divided into 3 groups: (1) chlorinated arylmethyl compds.
     containing at least 1 substituent C1 atom in the methyl group; (2)
     halogenated aliphatic hydrocarbons containing at least 1 CX3 group, where X is
     C1, Br, or I; (3) aliphatic compds. containing 1 CCl3 group attached
     to a strong polar group. The compds. tested include benzyl chloride,
     2,4-dichlorobenzyl chloride, 2,4,6-trichlorobenzyl chloride, benzal
     chloride, 2,6-dichlorobenzal chloride, benzotrichloride,
    o-chlorobenzotrichloride, 2,4-dichlorobenzotrichloride,
     1-trichloromethylnaphthalene, 1,1,1,3-tetrachloropropane,
     1,1,1-trichloropentane, 1,1,1-trichloroheptane, 1,1,1,9-tetrachlorononane,
     1,1,1,5-tetrachloropentane, 1,1,1,5-tetrachloro-2-pentene,
    octachloro-1,3-pentadiene, hexachloroethane, heptachloropropane,
    pentachloropropane, pentachloroethane, bromoform, iodoform, C
    tetrabromide, telomer mixture, trichloroacetic acid, Et trichloroacetate, Zn
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Vulcanization of GR-S with halogen compounds Sturgis, B. M.; Baum, A. A.; Trepagnier, J. H.

E. I. du Pont de Nemours & Co., Wilmington, DE

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trichloroacetate, Pb trichloroacetate, N-trichloroacetylurea, Na trichloroacetate, trichloromethanesulfochloride. All these compds. are unique in that they are primary vulcanizing agents for GR-S, yet do not vulcanize natural rubber in the absence of S. Group (1) members are, in general, the most active and they vulcanize GR-S at room temperature without any accessory agent. A large number of halogenated aromatic compds. including hexachlorobenzene, benzene hexachloride, 9,10-dichloroanthracene, 2,4,6-trichloromesitylene, benzotrifluoride, 2,4,6trichlorobenzotrifluoride, and other analogous F compds. did not vulcanize GR-S. They are highly activated by some metal oxides, particularly PbO; e.g., with the latter vulcanizates with 2.5% of o-chlorobenzotrichloride having phys. properties close to those of S vulcanizates are obtained. The oxides tested include CeO2, Cu20, CuO, Pb304, Pb02, Pb0, Mg0, Mg02, Mn02, Te02, Zn0, and Zn02. MgC03, CaC03, and lithopone also are activators. In group (2), F compds. are inactive, and when alone the C1 and Br compds. are inactive in the absence of a metal oxide; in fact, PbO is the only highly effective oxide. The following compds. were inactive or only slightly active: 1,2,3-tribromobutane, 1,2,3,4-tetrabromobutane, 1,2,3-tribromo-2methylpropane, 1,2,3,4-tetrachlorobutane, 1,2,3,3-tetrachlorobutane, 1,2,2,3-tetrachlorobutane, 2,2,3-tribromobutane, 1-4-dibromo-2-butene, 1,2-dibromo-1-chloroethane, 3,4-dibromohexane, 1,1,2,2-tetrabromoethane, hexachlorobutadiene, n-butyl iodide, 1,1,7-trichloroheptene, 1,3,3,4,5,6-hexachlorohexane, and methyl iodide. S retards the action of aliphatic halogenated compds. With group (3), both ZnO and S are activating agents and are necessary for the best quality of vulcanizates but they are not necessary to obtain vulcanizates of fair quality. PbO, MgO, Pb3O4, MoO3, and ZnO2 are less effective activators. That compds. of group (3) are primary vulcanizing agents with S present is evidenced by the fact that amines, e.g. tetraethylenepentamine, activate S alone but retard a combination of S and ethyl trichloroacetate. In general, the halogenated organic vulcanizing agents give GR-S vulcanizates which are characterized by particularly good resistance to elevated temps., e.g. aging at 100°. The mechanism of vulcanization by these compds. does not conform perfectly to any current theories of vulcanization by non-S compds., but the mechanism is possibly analogous to that of BzO2 (cf. Alfrey, Hendricks, Hershey, and Mark, C.A. 39, 5542.4), i.e.: initial decomposition into free radicals; reaction of the free radicals with the polymer, with elimination of an  $\alpha$ -methylenic H atom and formation of an odd electron; reaction of the polymer free radical with a polymer mol., probably with 1 of the double-bond electrons, with cross-linking; and continuation of the process, either by a sequence of cross-linking through the double bonds of addnl. chains or by chain transfer, until a termination reaction takes place.

**5324-68-5**, Mesitylene, 2,4,6-trichloro-

(as vulcanizing agent in Buna-S)

5324-68-5 CAPLUS

Benzene, 1,3,5-trichloro-2,4,6-trimethyl- (9CI) (CA INDEX NAME)

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